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1	1. A method to perform dual, sequential
2	diagnostic testing of the heart on the same
3	patient, with each half of the dual testing having
4	two parts, the first part being a baseline study and
5	the second part being the use of stress means
5	designed to exercise the heart during the second
7	part of the initial half of the dual test, and
3	immediately after its completion, de novo d-
9	ribose is administered for one hour or longer,
10	whereupon, the same two-part test is repeated as
11	the second half of the dual test.
l	2. The method of claim 1 in which from 12 to
2	60 grams or more of de novo d-ribose is
3	administered by mouth following completion of
4	the initial half.
1	3. The method of claim 1 in which stress can
2	be elicited by physical exercise to induce the
3	heart to contract more rapidly.
1	4. The method of claim 1 in which stress by
2	chemical inotropic means can be used to induce
3	the heart to contract more rapidly.
1	5. The method according to claim 4 in
2	which dobutamine is the chemical agent.

1	1	6. The method of claim 1 in which more than 60
2	2	grams of d-ribose are administered during and
3	3	after the test.
4	1	7. The method of claim 1 in which the various
5	2	stress scanning tests of the heart include but are
6	3	not limited to electrocardiographs,
7	4	echocardiographs, thallium scintigraphy, PET
8	5	(positron emission tomography) scanners, CT
9	6	(computerized tomography) scanners and MRI
10	7	(magnetic resonance imaging) scanners and
11	8	electron beam imaging scanners.
12	1	8. The method of claim 7 in which
13	2	electrocardiograph and sphygmotonograph
14	3	electrodes are attached to the patient and used for
15	4	monitoring purposes.
16	1	9. The method of claim 1 in which intravenous
17	2	infusion of d-ribose is used for at least one half
18	3	hour.
19	1	10. The method of claim 1 in which the heart
20	2	function having been improved diagnostically by
21	3	de novo d-ribose, said d-ribose is continued
22	4	therapeutically afterwards.
23	1	11. The method in which the minimum
24	2	practical levels of de novo d-ribose dosage is
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3	determined by serial imaging studies, each
4	following the other by more than 24 hours,
5	showing the degree of myocardial contractility
6	for a given dosage of d-ribose, for which any
7	non-invasive, immediately sequential imaging
8	procedure for the heart can be used.
1	12. The method of claim 7 in which the
2	determination of the heart rate and blood pressure
3	is done manually.
1	13. The method of claim 7 in which the Philips
2	Medical Systems' electrocardiographs are used
3	for the testing.
1	14. The method of claim 7 in which Holter
2	monitor means are used as conventionally used
3	on only one person for 24 to 48 hours.
1	15. The method of claim 13 in which the Holter
2	monitor is one of the Zymed 1810 family of
3	recorders using Windows.
1	16. The method of claim 13 in which said
2	scanning is done at fitness and health clubs.
1	17. The method of claim 1 in which when said
2	baseline scanning is reported as normal, the
3	baseline is repeated serially until an abnormality
4	occurs and then the d-ribose protocol followed.

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18. The method of claim 14 in which when the
Holter monitor is not used as conventionally
used, the software is written for intentional
sequential interrupting of the scanning so that
multiple individuals can be scanned on one unit
for recording, retrieval and storage over an
elapsed time period that could last up to 48 hours
of total although continually interrupted use.
19. The method according to claim 1 in which
the ribose part of the test is done first and the
baseline afterwards.